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Synthetic sea shell

Michael Rubner

The mechanical properties of natural substances such as bone and shell are envied by those involved in the fabrication of materials. A 'bricks-and-mortar' structure, assembled layer by layer, is the key to making sea shells.

For a materials scientist, cross-sectional images of the complex microstructures of naturally occurring hard materials such as bones and sea shells are awe-inspiring. Over many millions of years, nature has devised schemes to combine seemingly incompatible building-blocks — 'soft' organic proteins and 'hard' inorganic particles of calcium carbonate — in a manner that produces composite materials with the unusual combination of high strength, hardness and toughness. Imagine, however, that you could build such a structure as a mason would, one layer at a time, from the bottom up. Writing in *Nature Materials*, Tang and colleagues¹ explain how it can be done, using a molecular-level processing scheme known as layer-by-layer assembly^{2,3}.

Flexible soft materials that can undergo energy-absorbing molecular rearrangements during deformation are tough, but also very compliant. In contrast, rigid hard materials are stiff but often also very brittle, and they have little ability to absorb energy, so their toughness is low. To be strong, hard and tough, a material must be able to absorb a large amount of energy during mechanical deformation and also maintain high stiffness. In bone or shell, this desirable combination of properties is made possible by one key attribute — a bricks-and-mortar-like structure, made up of strongly interacting, nanometre-size building-blocks. The 'hard' bricks and 'soft' mortar are complementary in their response to stress and strain.

So far, attempts to mimic these structures with synthetic building-blocks have failed to produce a material with similarly impressive mechanical properties, because most conventional processing techniques simply do not offer the nanoscale level of control needed to create a highly regular bricks-and-mortar-type arrangement. Nature has no such difficulty with nanoengineering: it can assemble, in a regular manner, building-blocks of the right dimensions that interact strongly enough at their interfaces to allow the transfer of deformation energy between the rigid bricks and the softer mortar. Reproducing these elements synthetically is a challenge.

But this is exactly what Tang *et al.*¹ have achieved, through the alternating sequential deposition of negatively charged, nanometre-thick clay platelets (the bricks) and a positively charged polymer (the mortar).

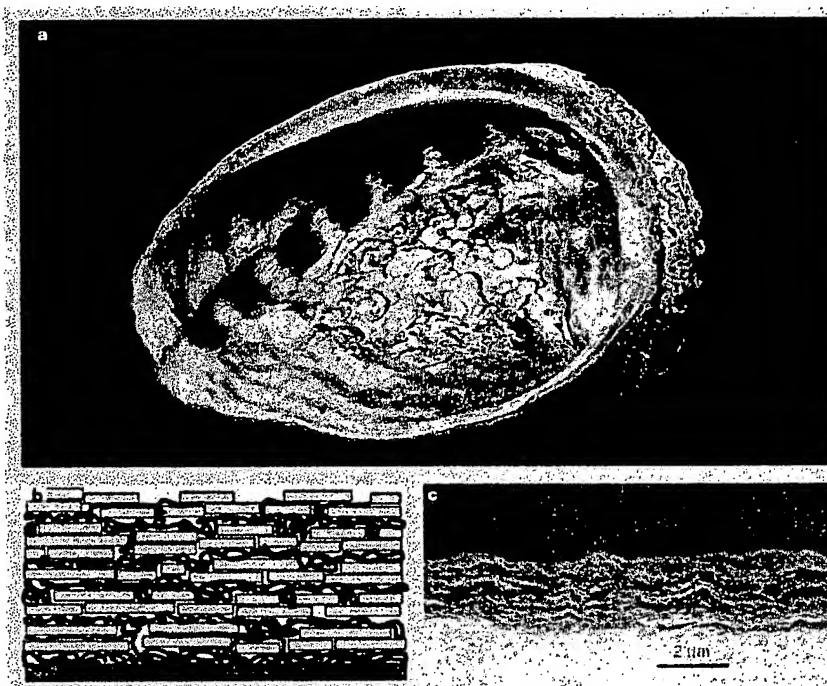


Figure 1 The 'bricks-and-mortar' approach. a, The natural strength, hardness and toughness of bone and shell are attributable to their nanoscale structure of calcium carbonate bricks and mortar-like protein layers. By mimicking this structure, Tang *et al.*¹ have created a new material with mechanical properties similar to nacre, or mother-of-pearl. b, Montmorillonite bricks (0.9 nm thick) are deposited layer by layer above a silicon-wafer substrate, alternating with polymer chains of mortar. c, Structures that are many layers deep can be built — this one has 100 brick and polymer layers — although the process is slow.

The primary driving force for this adsorption-based assembly, which is carried out entirely from dilute aqueous solutions of the materials, is electrostatic: the positively charged polymer chains are attracted to the negatively charged clay platelets, and vice versa. By assembling the clay platelets (in this case, a material called montmorillonite) and polymer chains one deposition step at a time, the authors are able to create a bricks-and-mortar-type arrangement that mimics the natural structure of nacre, the material known as mother-of-pearl (Fig. 1).

The polymer chains are arranged in coils and folds, physically pinned by relatively weak electrostatic interactions. As the material is deformed and the clay platelets begin to slide over each other, the polymer chains can undergo molecular rearrangements, through the breaking of 'sacrificial' ionic bonds and the concomitant unfolding of the coiled polymer chains. And this process, in turn, allows the material to absorb a lot of

deformation energy. The authors report that thin films assembled in this way have tensile strength and stiffness that approach those of seashell nacre. Their stiffness is significantly higher than the more disordered composites previously fabricated from similar materials using conventional methods.

As a step towards creating synthetic materials that truly mimic the mechanical behaviour of naturally occurring materials, this is an important advance. The true potential of this approach — to construct more complicated layered structures containing many types of building-blocks — has yet to be pursued. But, for example, this group has also shown that layer-by-layer assembled films containing carbon nanotubes have exceptional mechanical properties⁴. So this seems a promising way of fabricating multilayered, multi-component thin films, with molecular architectures designed to take full advantage of the complex interactions possible between many different types of materials. And as this

assembly process simply involves alternately dipping a substrate into dilute aqueous solutions of oppositely charged materials, it is easy to control the number, type and sequence of layers added to the film⁵.

There is a price to pay, however, for assembling the building-blocks one layer at a time. Such structures, particularly if they need to reach thicknesses in the micrometre range, require far longer fabrication times than more conventional processes such as spin-coating. Indeed, it is impressive that Tang *et al.*¹ have succeeded in creating free-standing, micrometre-thick films in this way (Fig. 1c).

Another challenge in working with charged polymers is that they are capable of adsorbing a lot of water, which in turn can degrade the mechanical properties by screening the ionic interactions that lend strength to the material. Tang *et al.* show that there is a significant decrease in mechanical performance when films are tested in high-humidity environments. But this problem

can be addressed, as many different types of materials can be assembled in these multilayer films, including those that are hydrophobic (water resistant) or that can be rendered hydrophobic by subsequent chemical or thermal treatments.

In any event, these model structures are sure to provide new insight into the behaviour of naturally occurring materials. Their applications could be widespread, from synthetic engineering of biological hard tissue to thin-film protective coatings.

Michael Rubner is in the Department of Materials Science and Engineering, Massachusetts Institute of Technology, 77 Massachusetts Avenue 13-5110, Cambridge, Massachusetts 02139, USA.
e-mail: rubner@mit.edu

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Telomeres

Taking the measure

Vicki Lundblad

Telomeres — the tips of chromosomes — need to be preserved, and this involves replenishing telomeric DNA when it has been eroded. But telomeres must not become too long, and one aspect of length control is now revealed.

In every organism, maintaining the integrity of the genome is a crucial endeavour. One aspect of genome maintenance involves protecting telomeres, the natural ends of linear chromosomes. This task is achieved by a suite of specialized protein complexes, which are anchored to chromosome ends through their association with further proteins that bind directly to telomeric DNA. The resulting structure prevents events that would be catastrophic for the genome, such as the loss of terminal DNA sequences or end-to-end chromosome fusions.

One of the complexes involved in telomere maintenance is an enzyme called telomerase, which adds DNA back to telomeres that have become eroded. Several other proteins also regulate this complex. But how the different proteins talk to one another — to keep telomeres the right length, to protect them, and to replicate them during cell division — is poorly understood. Writing on page 1013 of this issue and in *Current Biology*, respectively, Loayza and de Lange¹ and Colgin *et al.*² describe a crucial feature of the process by which telomerase can sense, and thus regulate, the length of individual chromosome ends.

Telomeric DNA is composed of G-rich repeats — reiterations of a short DNA sequence that does not code for protein and

is high in guanine (G) nucleic-acid bases. It also has a single-stranded stretch that overhangs the end of the double-stranded (duplex) telomeric region. This overhang is

the substrate for telomerase, which elongates chromosome ends by adding G-rich repeats. The importance of telomerase is evident from studies of yeast and human cells in which reductions in telomerase levels produce a steady decline in telomere length that eventually blocks cell division. Not surprisingly, then, telomerase is highly active in systems such as the blood and reproductive system, which rely on continuous replenishment through cell proliferation³. Much to the interest of cancer biologists, telomerase levels are also increased in most human tumours, providing a potential target for the development of anticancer drugs⁴.

In normal cells, telomerase activity is carefully controlled by several mechanisms. For instance, subunits that are part of the telomerase complex itself can positively regulate the enzyme, for example by mediating recruitment of the complex to chromosome ends⁵. Surprisingly, proteins that bind to the duplex region of the telomere can also be potent regulators, even though they do not appear to associate physically with telomerase. These duplex-binding proteins — which include Rap1 in budding yeast and the TRF1 and TRF2 proteins in human cells — can 'count' the number of G-rich repeats and, when telomeres become overly long, inhibit further telomerase activity^{6,7}.

Missing from this elegant proposal for telomere-length regulation, however, is an explanation for how information from the duplex portion of the telomere is relayed to the very tip of the chromosome — the site of telomerase action. To address this, Loayza and de Lange¹ and Colgin *et al.*² turned to a recently discovered human protein called

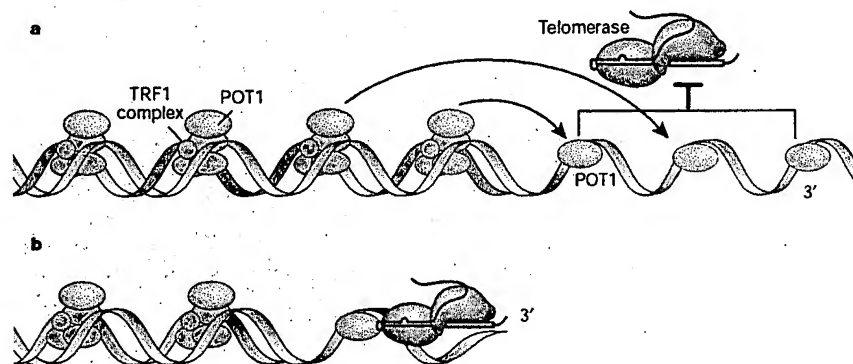


Figure 1 Model for telomere length control. Telomeres are found at the end of linear chromosomes, and their length must be precisely regulated — a process that involves the POT1 protein in human cells^{1,2}. Loayza and de Lange¹ have shown that POT1 binds to the TRF1 complex on the double-stranded (duplex) portion of telomeres. TRF1 complexes sense the length of the telomere, and the authors propose that this information is transmitted to telomerase (an enzyme that extends telomeres) via POT1, by transferring POT1 to the single-stranded overhang at the telomere tip. **a**, When the telomere is long enough, the levels of POT1 on the overhang are high, and telomerase is inhibited. **b**, When the telomere is too short, little or no POT1 is transferred to the end, and telomerase is no longer inhibited, allowing it to add DNA back to the telomere. Colgin *et al.*² have proposed that POT1 may also act as a positive regulator of telomerase when present at the single-stranded terminus. It might do so via a direct interaction with telomerase, in an analogous way to how the yeast Cdc13 protein regulates telomerase^{5,11}.

Cooperative Formation of Inorganic-Organic Interfaces in the Synthesis of Silicate Mesostructures

Monnier, F. Schüth, Q. Huo, D. Kumar, D. Margolese, S. Maxwell, G. D. Stucky,* M. Krishnamurty, P. Petroff, A. Firouzi, M. Janicke, B. F. Chmelka

is presented to explain the formation and morphologies of surfactant-silicate mesostructures. Three processes are identified: multidentate binding of silicate oligomers to a cationic surfactant, preferential silicate polymerization in the interface region, and charge density matching between the surfactant and the silicate. The model explains experimental data, including the transformation between lamellar and hexagonal mesophases, and provides a guide for predicting conditions that favor the formation of hexagonal, or cubic mesostructures. Model Q²³⁰ proposed by Mariani and his co-workers satisfactorily fits the x-ray data collected on the cubic mesostructure material. The model suggests that the silicate polymer forms a unique infinite silicate sheet sitting on a minimal surface and separating the surfactant molecules into two disconnected volumes.

vention of a new family of mesoporous materials, designated M41S, by scientists at Mobil Oil Corporation (1), has dramatically expanded the range of crystallographically defined pore sizes from the micro- (3 Å) to the mesopore (20 to 100 Å). The synthesis uses ordered arrays of surfactant molecules as a "template" for the in situ polymerization of silicates. Mesoporous materials obtained by this method exhibit several remarkable features: (i) a wide range of pore sizes and shape, as compared to microporous materials; (ii) fine adjustment of the pore size within the limits stated above; (iii) high thermal and hydrolytic stabilities; and (iv) a very high degree of pore ordering over micrometer distances. These unusual properties are a result of the interplay between organization of the surfactant molecules and silicate species in the aqueous phase.

Stucky et al. (2) outlined two general models for the formation of the mesophases. The first model assumes that the primary structure-directing element is the surfactant liquid crystal phase. The second model suggests that the addition

of the silicate orders the subsequent silicate-encased surfactant micelles. These general models, however, are insufficient for establishing the mechanistic understanding needed for better control of the synthesis process, which is key to efforts aimed at improving or adding to this exciting new class of materials. On the basis of experimental results, we present here a more detailed model of the mesophase formation process, which explains presently known experimental data and successfully predicts conditions needed for the synthesis of desired structures. We believe that this model can be generalized to the synthesis of other nonsiliceous materials as well.

From considerations in surfactant and silicate chemistry, three closely coupled phenomena are identified as crucial to the formation of surfactant-silicate mesophases. These include: (i) multidentate binding of silicate oligomers, (ii) preferred polymerization of silicates at the surfactant-silicate interface, and (iii) charge density matching across the interface.

Mesostructure syntheses can be carried out under conditions in which the silicate alone would not condense (at pHs from 12 to 14 and silicate concentrations of 0.5 to 5%) and the surfactant cetyltrimethylammonium (CTA⁺) alone would not form a liquid crystal phase. In fact, surfactant-silicate mesophases can form at surfactant concentrations as low as 1%, a regime in the CTABr-water phase diagram in which only micelles are present. For a CTABr-water solution at typical surfactant-silicate synthesis temperatures in the absence of silicates, a hexagonal phase is favored at surfactant concentrations from ~25 to 70% by weight whereas a lamellar phase forms at concentrations above 70% (3, 4). Never-

theless, a solid mesophase precipitate is formed, the structure of which will be discussed below, as soon as surfactant (chain length of 8 to 20 carbon atoms) and silicate solutions are combined. The rapidity of this precipitation indicates that there is a strong interaction between the cationic surfactant and anionic silicate species in the formation of surfactant-silicate mesophases.

We performed syntheses aimed at identifying conditions important for the formation of mesoporous materials over a wide range of reactant compositions and temperatures (5). For the purpose of investigation, we found that we could slow the evolution of the surfactant-silicate systems by undertaking the syntheses at moderate temperatures (between 30° and 100°C) (6). During freeze-dry kinetic experiments with CTACl used as the surfactant, a layered (lamellar) material with a primary *d* spacing (repeat distance) of 31(±1) Å was produced, together with amorphous silica, after reaction times on the order of 1 min. For the synthesis conditions given in Fig. 1, the lamellar mesophase disappears after approximately 20 min, at which point the diffraction pattern of the hexagonal mesostructure is simultaneously detected. This hexagonal material has a primary *d* spacing of 40(±1) Å and attains its final degree of ordering after ~10 hours (7).

A layered material with a primary *d* spacing of 31(±1) Å (Fig. 2, pattern A) can be isolated in pure form (8); a transmission electron microscopy (TEM) micrograph of this mesostructure is depicted in Fig. 3. The variation of the *d* spacing as a function of the chain length of a cationic surfactant C_nH_{2n+1}[N(CH₃)₃]⁺ (for 14 ≤ *n* ≤ 22) is 1.0 to 1.2 Å per carbon, which

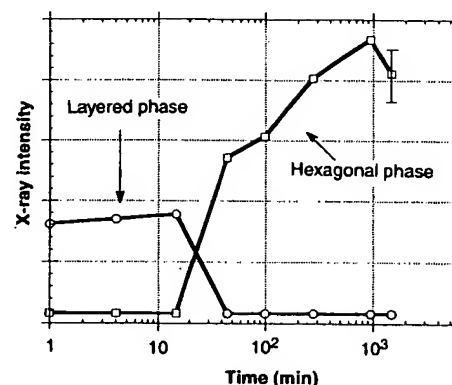


Fig. 1. Time evolution of the intensity of x-ray diffraction features associated with layered and hexagonal (M41S) mesostructures at 348 K. The layered material is precipitated rapidly, whereas the hexagonal material appears later, as a result of a higher degree of silica polymerization. The composition of the reaction mixture was as follows: 1 M SiO₂:0.025 M Al₂O₃:0.115 M Na₂O:0.233 M CTACl:0.089 M TMAOH:125 M H₂O.

* Department of Chemistry, University of California, Santa Barbara, CA 93106, and Department of Physique Sciences II, 1211 Geneva, Switzerland.

† Department of Chemistry, University of California, Santa Barbara, CA 93106, and Institut für Chemie, Johannes-Gutenberg-Universität, Mainz, Germany.

‡ D. Kumar, D. Margolese, R. S. Maxwell, G. D. Stucky, Department of Chemistry, University of California, Santa Barbara, CA 93106.

§ P. Petroff, Materials Department, University of California, Santa Barbara, CA 93106.

|| M. Janicke, B. F. Chmelka, Department of Nuclear Engineering, University of California, Santa Barbara, CA 93106.

Correspondence should be addressed to G. D. Stucky.

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corresponds to a monolayer assembly. If this new layered material is hydrothermally treated at 373 K (pH = 7), it is converted to the hexagonal mesostructure over 10 days, with intermediate and final x-ray patterns shown in Fig. 2, patterns B and C, respectively. During this transformation the degree of silica polymerization increases, as measured by the relative number of incompletely condensed (Q^3) and fully condensed (Q^4) silicon atoms determined by ^{29}Si magic-angle spinning nuclear magnetic resonance spectroscopy. The ratio between Q^3 and Q^4 silicon decreases from typical values of 1.0 for the layered material to 0.4 to 0.55 for the hexagonal mesostructure, reflecting a significant increase in the number of silicon atoms fully coordinated to other silicate nearest neighbors.

Mesophase formation and associated silica polymerization are intimately tied to Coulombic interactions between surfactant and silicate species at the micelle interfaces. Silicates present in the form of monovalent monomers, $\text{Si}(\text{OH})_3\text{O}^-$, however, are expected to have little energetic advantage over other monovalent anions competing for access to the cationic surfactant head groups. At high pH, the reaction mixture also contains small silica oligomers (three to seven silicon atoms) of varying degrees of polymerization and charge (9). These oligomers are appreciably more acidic ($\text{p}K_a \sim 6.5$) than the monomer or dimer species [$\text{p}K_a$ 9.8 and 10.7, respectively (10)], although all such silicates will be highly dissociated under the high pH conditions used here (11).

The oligomeric silica polyanions, however, can easily act as multidentate ligands for the cationic head groups of the surfactant, leading to a strongly interacting surfactant-silicate interface. Indeed, the interaction of ionic surfactants with polyions of opposite charge encourages strong cooperative binding, manifested by increases in the binding constants of up to two orders of

magnitude in similar systems (12). Preferential multidentate binding of the silicate polyanions causes the interface to quickly become populated by tightly held silicate oligomers, which can subsequently polymerize further. Silicate polymerization within the surfactant-silicate interface region is favorable for two related reasons: (i) the concentration of silicate species near the interface is high and (ii) their negative charges are partially screened by the surfactant. Furthermore, as polymerization proceeds, the formation of highly connected silicate polyanions, which act as very large multidentate ligands, further enhances the cooperative binding between the surfactant and silicate species.

Multidentate ionic binding in surfactant-silicate systems has an important consequence; namely, it leads to precipitation of a given mesophase from solution. Through the interactions driving the precipitation process, the appearance of a given mesostructure is established, although this process is expected to operate on a different time scale from polymerization of the silica, which accounts ultimately for the thermal, mechanical, and hydrolytic stability of the final material. If small silica oligomers are present in sufficient quantity, precipitation of the surfactant-silicate system is primarily the result of electrostatic interactions, combined with packing constraints associated with the hydrophobic surfactant chains. Whereas precipitation is fast and essentially thermodynamically controlled, silica polymerization into a strong and extended framework is slow and reaction rate-limited. This two-stage process is in agreement with experimental findings that contrast the mesostructures obtained at room temperature after short reaction times with those obtained at high temperature after long reaction times: very similar x-ray patterns are obtained for both sets of conditions, indicating identical precipitated mesostructures; however, the materials syn-

thesized by the low-temperature route are thermally and mechanically much less stable than the high-temperature analogs. The coupling between the precipitation and polymerization processes in surfactant-silicate systems provides the basis for the lamellar-to-hexagonal mesophase transformation in a way that we now describe.

The resemblance, in shape and size, of the surfactant-silicate mesostructures with the corresponding water-surfactant liquid crystal phases indicates that the interactions responsible for these morphologies are of a similar nature. The governing role of the head-group area (A) in the selection of a particular mesophase has already been recognized in water-surfactant systems: the favored mesophase is that which permits A to be closest to its optimal value A_0 , while maintaining favorable packing of the hydrophobic surfactant chains (13). In surfactant-silicate systems, the value of A_0 is strongly affected by electrostatic and steric interactions between the silicate and surfactant micelle species. More quantitatively, its value is obtained by minimizing the Gibbs free energy G as a function of A

$$A_0 \rightarrow (\partial G / \partial A) = 0 \quad (1)$$

$$G(A, \rho) = G_{\text{intra}}(A) + G_{\text{wall}}(\rho) + G_{\text{inter}}(A, \rho) + G_{\text{sol}} \quad (2)$$

where G_{intra} reflects the van der Waals forces and conformational energy of the hydrocarbon chains and the van der Waals and electrostatic interactions of the head group within a single micelle; G_{wall} accounts for the polysilicate structural free energy, including the solvent, counterion, and silicate van der Waals and electrostatic interactions within the inorganic silicate framework or "wall"; G_{inter} reflects the van der Waals and electrostatic effects associated with wall-micelle and micelle-micelle interactions; G_{sol} describes the solution phase; and ρ specifies the composition of the various species within the wall. The chemical potentials of these species are set by the concentration of the corresponding species in the aqueous solution, as accounted for by G_{sol} .

Fig. 2. Powder x-ray diffraction patterns of surfactant-silicate mesostructures precipitated from the same reaction mixture (1 M SiO_2 ; 0.034 M Al_2O_3 ; 0.07 M Na_2O ; 0.27 M CTAB; 0.14 M TMAOH; 0.28 M TMB; 100 M H_2O), and then treated hydrothermally at 373 K for different times. X-ray patterns are shown for (curve A) the initially precipitated layered material, (curve B) an intermediate material, and (curve C) the M41S hexagonal mesostructure acquired 0, 1, and 10 days, respectively, after initiation of the hydrothermal treatment.

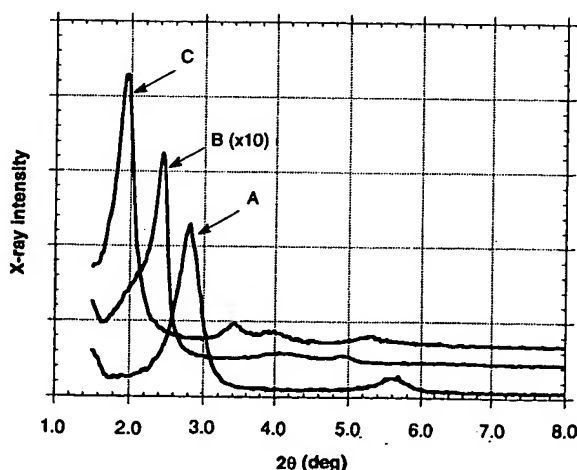


Fig. 3. Transmission electron micrograph of the layered surfactant-silicate mesostructure whose x-ray data are shown in Fig. 2 (curve A). The d spacing of this material is $31(\pm 1)$ Å.

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ature route much less than the other analogs. The precipitation and surfactant-silicate interaction for the phase transformation describe. The term G_{wall} drives the chemistry within the wall, including the polymerization process, and contains the structural constraints responsible for the multidentate binding. The term G_{inter} establishes the interaction between A and the state of the wall described by ρ . This coupling across the interface can be understood in terms of electrostatic interactions (which most likely predominate), whereby the silicate charge density within the wall, ρ_e , is mutually screened by the charges on the surfactant head groups, which have an average face charge density of $1/A$. Thus, the electrostatic interactions link A_0 , as defined in Eq. 1, with ρ_e , a relation we refer to as "charge density matching." Such interdependent electrostatic effects control the degrees of surfactant intercalates in different mica-type silicates (15) and have been invoked to explain the "self-replication" process of silica layers in purely inorganic systems (16).

In surfactant-silicate systems, polymerization driven by G_{wall} will profoundly affect the mechanism providing a mechanism to explain the transition between the lamellar and hexagonal

mesophases. In the early stage of the synthesis, the presence of highly charged silica oligomers favors a small value of A_0 , which can be achieved with a lamellar surfactant configuration. As rearrangement and polymerization of the silicate species proceed, the density of anionic silanol groups diminishes, so that A_0 increases, while the number of compensating cations decreases. At the same time, the wall thickness can decrease from its initial value without energy cost, because the most stable ionized silanol groups are confined to the wall surface, thus reducing repulsive dipole-dipole interactions between the two opposite-facing wall surfaces. The silicate wall is still poorly condensed during early stages of the synthesis, allowing the system to increase its A toward A_0 by adopting the hexagonal structure according to charge-density matching criteria. Under these circumstances, the wall thickness simultaneously decreases to keep the volume ratio CTA/SiO₂ constant. The actual wall thickness has been estimated to be 10 to 11 Å (17) for the lamellar mesophase and 8 to 9 Å (18) for the hexagonal mesophase. Simple geometrical arguments can be used to show that these values are consistent with a constant CTA/SiO₂ volume ratio throughout the phase transition.

The regularity of the product mesostructures supports mediation of the silicate wall

thickness during the assembly process. The high efficiency of this mediation is reflected by the experimental observation that the wall thickness of the hexagonal phase is essentially constant (8 to 9 Å) over a wide range of reaction conditions, independent of the surfactant chain length, and by the clearly hexagonal, as opposed to circular, pore shape established by both high-resolution TEM and modeling of the powder x-ray diffraction patterns (19). Control of the silicate wall thickness is undoubtedly related to the double layer potential: silicate species are only accumulated at the surfactant interface to the extent necessary for charge compensation. Polymerization normal to the interface, which would thicken the wall or produce amorphous bulk SiO₂, does not occur because of the strong electrostatic repulsion produced by the high negative charge on the silicate species at pH 12 and above (10).

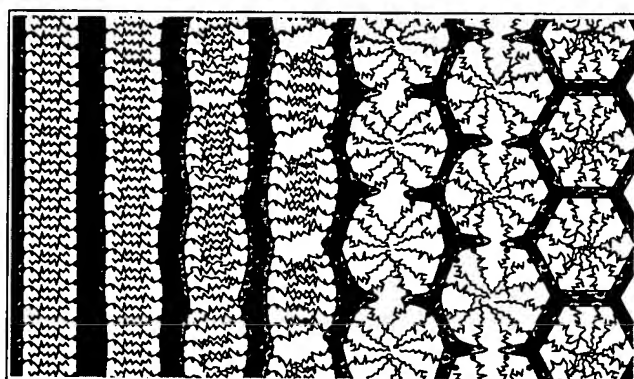
Figure 4 shows a mechanism consistent with current experimental investigations by which the lamellar-to-hexagonal mesophase transformation may occur. Silica polymerization leads to an increase in interfacial area that is achieved through corrugation of the lamellar surfactant-silicate sheets. As implied in the final step, this corrugation progresses until connection between the sheets is made at the cusps, resulting ultimately in the formation of the hexagonal mesophase. Another way to accommodate the change in A would be to maintain a planar structure while tilting the hydrocarbon chains. Such a transition, however, is entropically disfavored by the restrictive chain configuration this suggests.

Yanagisawa *et al.* (20) recently reported a hexagonal mesostructure, with pore dimensions similar to that of M41S, produced by the inclusion of CTA⁺ cations into the sheet silicate kanemite. During their synthesis these researchers observed a layered intermediate that subsequently transformed into a hexagonal phase material. This process is probably driven by the same forces as the transformation we report, although it is not yet clear to what extent the kanemite structure is preserved during the conversion. If the pH is sufficiently basic, for example, the sheets can be partially or fully destroyed during the process.

We propose that the surfactant-silicate mesophase structure is governed primarily by the terms G_{intra} and G_{inter} of Eq. 2. In this respect, the main effect of the silicate wall and of the reaction conditions are to determine A_0 . This provides predictive capability for establishing the reaction conditions that favor the lamellar or the hexagonal mesophases. We have tested this model experimentally by monitoring the effects of pH and the degree of polymerization of the silica source on the mesostructure syntheses, with the results summarized in Fig.

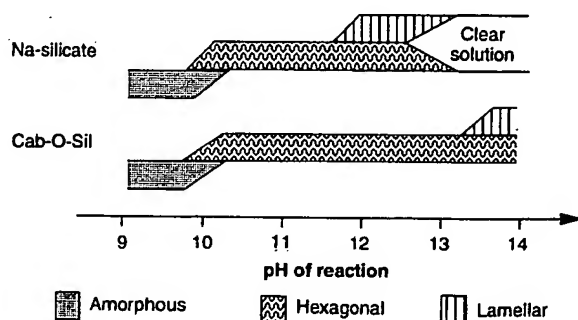
Fig. 4. Schematic diagram

of the mechanism proposed for the transformation of a surfactant-silicate system from the lamellar to the hexagonal mesophase. On the left, small silica oligomers (not shown explicitly in the gray SiO₂ regions) act as multidentate surfactants, which have sufficiently high charge density to maintain a lamellar surfactant configuration. As polymerization of the silica proceeds, the diminished charge density of larger silica oligomers increases the average head-group area of the surfactant assembly, driving the transformation into the hexagonal mesophase.



■ SiO₂ ——— Reaction coordinate →

Fig. 5. Chart showing the approximate domains of formation of the lamellar and hexagonal surfactant-silicate mesophases, as functions of surfactant and silica source. Cab-O-Sil is composed of ~100 Å oligomeric particles, whereas Na-silicate is composed of hydrolyzed and essentially monomeric silicates.



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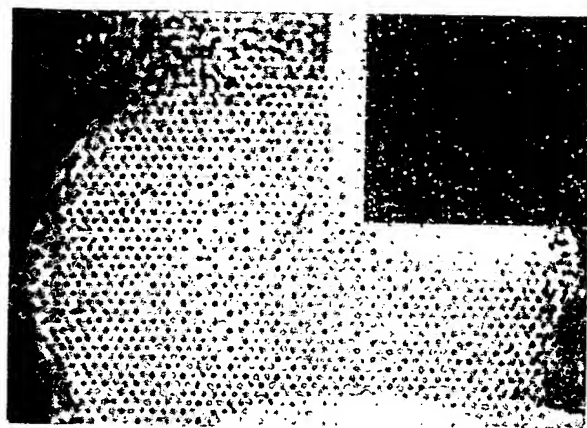
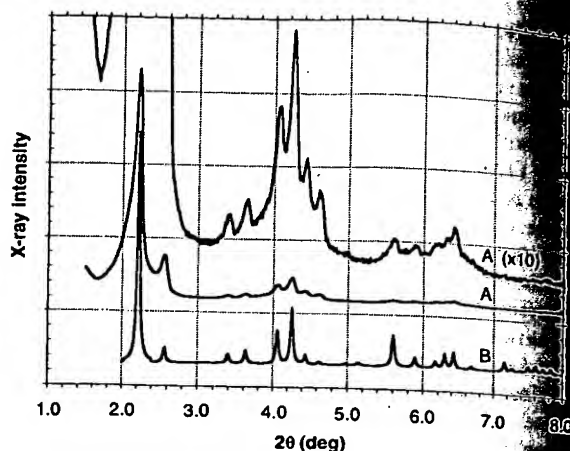


Fig. 6 (left). Transmission electron micrograph of the cubic surfactant-silicate mesostructure showing an ordered ~ 2000 Å aggregate viewed along its [111] axis. **Fig. 7 (right).** X-ray powder diffraction pattern of the cubic mesostructure, with $Ia\bar{3}d$ symmetry, synthesized from a reac-



tion mixture with a molar composition of 1 M TEOS:0.25 M Na₂O:0.63 M CTACl:62 M H₂O for 3 days at 373 K (curve A). Calculated diffraction pattern using the Q^{230} model proposed by Mariani *et al.* (21) with a lattice parameter $a = 97.3$ Å (curve B).

5. These results lead to the following conclusions in accordance with our predictions:

1) The lamellar phase is favored at high pH and for a low degree of polymerization of the silica source.

2) The hexagonal phase is favored at low pH and for a highly polymerized silica source.

In addition, we investigated the influence of the ionic strength on the surfactant-silicate assembly process by performing the synthesis in a reaction solution also containing 1 M NaCl. The presence of the salt decreased the regularity of the material, as reflected by a reduction in the number of peaks in the x-ray pattern (from four to two). This effect, expected only at high ionic strengths, is attributed to perturbation of the double layer potential. The strong binding constant of silicate species compared to other ions makes this effect negligible at lower ionic strengths and explains why mesostructure syntheses are relatively insensitive to other counterions in the reaction mixture.

The existence of the cubic mesophase described by Beck *et al.* (2) is strongly supportive of the important role of $G_{\text{intra}} + G_{\text{inter}}$ in the formation of surfactant-silicate mesostructures. Indeed, there is remarkable similarity between the cubic mesophase, which we have recently synthesized and characterized, and the $Ia\bar{3}d$ phase found in the water-CTAB system (4). A TEM image of the cubic mesostructure material (Fig. 6) shows an ordered ~ 2000 Å aggregate. The x-ray powder spectrum (Fig. 7) agrees very well with the model Q^{230} proposed by Mariani *et al.* (21) for water-surfactant systems. For this structure, it is appealing to conjecture that the midplane of the silicate wall sits on a gyroid periodic minimal surface (22). Such a structure can then be viewed as a single infinite silicate

sheet that separates the surfactant species into two equal and disconnected volumes. This so-called bicontinuous phase will be formed when the value of A_0 set by the reaction conditions is close to the value of the $Ia\bar{3}d$ phase, namely, when pH and the CTA/SiO₂ ratio are high. It is advantageous for the silicate wall to occupy a periodic minimal surface, because it can maximize the wall thickness for a given CTA/SiO₂ volume fraction.

The leading role of $G_{\text{intra}} + G_{\text{inter}}$ in directing mesostructure formation provides a foundation for identifying potential replacement candidates for silicon in the synthesis of mesoporous inorganic frameworks. The principal criteria are that the inorganic component must be capable of forming flexible polyionic species, that extensive polymerization of the inorganic component must be possible, and that charge density matching between the surfactant and inorganic species has to occur. In other words, when G_{sol} plays a benign role, G_{wall} must not dominate $G_{\text{intra}} + G_{\text{inter}}$ in order that the mesostructure form.

In addition to binding efficiently to the surfactant interface, the best inorganic candidates will have a tendency to form glasses easily. Silicates are certainly a prototypic system in view of the ease with which they form oligomeric anions with varying degrees of polymerization. Other systems, however, may also fulfill these requirements, including transition metals, such as vanadium, or main group elements, such as boron, which can form polyanions and condense. One can also speculate about a reversed system in which an anionic surfactant is used to precipitate a cationic metal oxide precursor, the laurylsulfate-iron oxide system representing one candidate example.

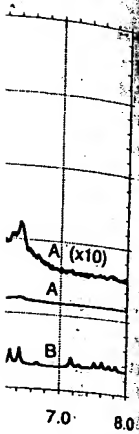
Existing experimental data thus far con-

firm the trends predicted for the formation of surfactant-silicate mesostructures by the qualitative model outlined above. Cooperative binding provides an explanation for the strong interactions needed to precipitate mesophases from dilute solutions. Preferential polymerization of silicates in the region of the interface together with a double layer control of the wall thickness are responsible for the high regularity of the surfactant-silicate mesostructures. Charge density matching establishes a link between the chemical composition and structure of the silicate wall and the formation of a particular mesostructure. We expect that these perspectives will stimulate and guide experiments aimed at producing and exploiting a better understanding of this exciting class of materials.

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expression "surfactant-silicate" is used here
a comprehensive term for materials syn-
thesized using a mixture of surfactant and silica
regardless of the particular structure.
transformation between the lamellar and
hexagonal mesophases was observed after
drying, as well as air-drying, the filtered
samples.
Addition of trimethylbenzene (TMB) to the reac-
tion mixture stabilizes the lamellar mesophase.
Experiments have shown that the 31(±1) Å re-
peat distance for the layered material shown in
Figs. 2A and 3 is preserved over a range of TMB
concentrations between 0.5 and 3.0 M, whereas
at lower TMB concentrations the hexagonal
mesostructure is the favored product. Stabiliza-
tion of the lamellar mesophase likely occurs
because TMB dissolved within the surfactant
hydrocarbon assemblies contributes to the hy-
drophobic chain volume. This increase in surfac-
tant chain volume increases the value of A_0 at
which the lamellar-to-hexagonal mesophase
transformation occurs, according to a simple
geometric model (13). Thus, the mesostructural
transformation depicted in Fig. 2 is a conse-
quence of hydrothermal removal of TMB from
within the surfactant chain assembly, combined
with an increase in A_0 . This conclusion is sup-
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dimensionless parameter $g = V/A_0\ell_c$ as a means
of determining the preferred configuration of a
surfactant assembly, where V is the volume of
the hydrophobic chain and ℓ_c is the characteris-
tic chain length. According to this treatment,
cylindrical micelles will form if $g < 1/3$, cylindrical
micelles if $1/3 < g < 1/2$, vesicles or bilayers if
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head-group diameter.
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An Unnatural Biopolymer

Charles Y. Cho, Edmund J. Moran,* Sara R. Cherry,
James C. Stephans, Stephen P. A. Fodor, Cynthia L. Adams,
Arathi Sundaram, Jeffrey W. Jacobs, Peter G. Schultz†

A highly efficient method has been developed for the solid-phase synthesis of an "unnatural biopolymer" consisting of chiral aminocarbonate monomers linked via a carbamate backbone. Oligocarbamates were synthesized from *N*-protected *p*-nitrophenyl carbonate monomers, substituted with a variety of side chains, with greater than 99 percent overall coupling efficiencies per step. A spatially defined library of oligocarbamates was generated by using photochemical methods and screened for binding affinity to a monoclonal antibody. A number of high-affinity ligands were then synthesized and analyzed in solution with respect to their inhibition concentration values, water/octanol partitioning coefficients, and proteolytic stability. These and other unnatural polymers may provide new frameworks for drug development and for testing theories of protein and peptide folding and structure.

Polypeptides have been the focus of considerable attention with respect to their structure and folding, biological function, and therapeutic potential. The development of efficient solid-phase methodology for the synthesis of peptides (1), peptide derivatives (2), and large peptide libraries (3–8) has greatly facilitated these studies. The development of efficient methods for the synthesis of unnatural biopolymers (9–11) composed of building blocks other than amino acids may provide new frameworks for generating macromolecules with novel properties. For example, polymers with improved pharmacokinetic properties (such as membrane permeability and biological stability) might facilitate drug discovery, and polymers with altered conformational or hydrogen-bonding properties may provide increased insight into biomolecular structure and folding. We report the highly efficient solid-phase synthesis of oligocarbamate polymers from a pool of chiral aminocarbonates and the synthesis and screening of a library of oligocarbamates for their ability to bind a monoclonal antibody (mAb).

The oligocarbamate backbone (Fig. 1), in contrast to that of peptides, consists of a chiral ethylene backbone linked through relatively rigid carbamate groups. The α carbon, like that of peptides, is substituted

with side chains that contain a variety of functional groups. Although the β carbon is unsubstituted in our initial target, additional backbone modifications (and conformational restriction) can be incorporated via alkylation of the β carbon or the carbamyl nitrogen. Oligocarbamates were synthesized from a pool of optically active *N*-protected aminocarbonates (Fig. 2) which, in turn, were derived from the corresponding optically active amino alcohols. The latter are either commercially available or can be prepared in chiral form by reduction of the *N*-hydroxysuccinimide or pentafluorophenyl esters of *N*-protected amino acids (12). The α -amino group was protected with the use of either nitroveratryl chloroformate (13) (NVOC-Cl) (for photochemical deprotection) or fluorenylmethyl-*N*-hydroxysuccinimide carbonate (Fmoc-OSu) (for base-catalyzed deprotection) (14). When necessary, side chains were protected as acid-labile *tert*-butyl esters, ethers, or carbamates. Protected amino alcohols were converted to the corresponding *N*-protected *p*-nitrophenyl carbonate monomers by reaction with *p*-nitrophenyl chloroformate in pyridine/CH₂Cl₂, generally in >80% yield. The carbonate monomers are stable for months at room temperature.

Solid-phase synthesis of oligocarbamates involves the sequential base-catalyzed or light-dependent deprotection of the α -amino group of the growing polymer chain followed by coupling to the next protected *p*-nitrophenyl carbonate monomer (Fig. 2). The *N*-protected "hydroxy-terminal" residue was covalently attached to polystyrene resin containing either *N*-protected *p*-alkoxybenzyl amino

C. Y. Cho, E. J. Moran, S. R. Cherry, J. C. Stephans, P. G. Schultz, Department of Chemistry, University of California, Berkeley, Berkeley, CA 94720.
S. P. A. Fodor, C. L. Adams, A. Sundaram, J. W. Jacobs, Affymax Research Institute, 4001 Miranda Avenue, Palo Alto, CA 94304.

*Present address: Ontogen Corporation, 2325 Camino Vida Roble, Carlsbad, CA 92009.

†To whom correspondence should be addressed.

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most weakly coupled QDs is as low as the spectral resolution of our spectrometer, 40 μeV , a value comparable to the smallest linewidth reported so far (6). The decrease of the exciton lifetime for the antibonding state with increasing level separation can be described in a simple two-level picture involving acoustical phonon scattering from the upper state into the lower state (19). This would lead to a cubic dependence of this linewidth on the energy level separation, which is also observed experimentally.

CEO has proven to be a versatile method for the fabrication of zero-dimensional objects of well-defined size, shape, and position. The excellent optical quality, manifested in extremely narrow emission lines, and the high degree of homogeneity accessible with this method permit the precise tailoring of the quantum-mechanical coupling between these nanoscale structures. As an extension to this work, we propose the use of higher barriers, narrower QWs, and the incorporation of indium into the wells. All these measures should increase the binding energy of excitons to the QDs. The use of strained InGaAs QWs, in particular, is expected to enhance this binding energy drastically because the strain can be almost completely elastically relaxed at the intersections. This might open a route to experimental investigation of a variety of quantum mechanics textbook examples previously inaccessible by other means, such as going from artificial atoms to molecules to an artificial one-dimensional solid.

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same objective lens and directed to a confocal imaging system, which defined a nearly diffraction-limited detection range of FWHM = 800 nm. The PL signal was dispersed with a triple-grating spectrometer (spectral resolution = 40 μeV) and detected with a cooled charge-coupled-device camera.

16. The PL energy of (110) QWs is red-shifted with respect to that of (001) QWs of identical thickness because of different heavy hole masses in these two directions.
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Organically Modified Aluminosilicate Mesostructures from Block Copolymer Phases

Markus Templin, Achim Franck, Alexander Du Chesne, Heike Leist, Yuanming Zhang, Ralph Ulrich, Volker Schädler, Ulrich Wiesner*

Organically modified aluminosilicate mesostructures were synthesized from two metal alkoxides with the use of poly(isoprene-*b*-ethyleneoxide) block copolymers (PI-*b*-PEO) as the structure-directing molecules. By increasing the fraction of the inorganic precursors with respect to the polymer, morphologies expected from the phase diagrams of diblock copolymers were obtained. The length scale of the microstructures and the state of alignment were varied using concepts known from the study of block copolymers. These results suggest that the use of higher molecular weight block copolymer mesophases instead of conventional low-molecular weight surfactants may provide a simple, easily controlled pathway for the preparation of various silica-type mesostructures that extends the accessible length scale of these structures by about an order of magnitude.

Currently, a great deal of attention is being paid to the synthesis of complex inorganic materials with long-range order (1). Such materials could find applications in catalysis, membrane and separation technology, and molecular engineering (2). A typical approach is the use of organic structures formed through self-assembly as structure-directing agents. The final morphology is then determined by the cooperative organization of inorganic and organic molecular species into three-dimensionally structured arrays, a concept also discussed in the context of biomineralization (3). This strategy has already been successfully used in the preparation of inorganic mesoporous materials (4). Different pathways, where the driving forces of the cooperative organization are either ionic (5) or based on hydrogen bonds (6), have been described in vastly different concentration regimes (7). Pore sizes of 20 to 100 Å are commonly obtained in this way.

Here, we used block copolymers of high-

er molecular weight to make the transition from the small to the large mesoscopic regime (up to several tens of nanometers) of silica-type mesostructures. Bagshaw *et al.* previously used block-type low-molecular weight surfactants as templating agents to produce mesoporous molecular sieves (6). Higher molecular weight block copolymers have been used to stabilize inorganic metal or semiconductor nanoparticles (8). However, they all produce solid particles with morphologies never very far from spherical (9). An example of a different shape of inorganic material in a random-coil organic homopolymer is the synthesis of randomly distributed inorganic nanowires (10). Most recently, block copolymers have been used to control the growth of anisotropic inorganic crystals (11).

Block copolymer materials are similar to low-molecular weight nonionic surfactant solutions with respect to their general phase behavior (12). The phase diagrams of these materials have been elucidated by numerous experimental and theoretical studies (13). The combination of inorganic siliceous components in a hybrid material with block copolymers is appealing for various reasons.

Max-Planck-Institut für Polymerforschung, Postfach 3148, 55021 Mainz, Germany.

*To whom correspondence should be addressed.

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First, a blend of desirable macroscopic properties (mechanical, thermal, and so forth) in the final product can be expected. Because the block copolymer chemistry (architecture, chain length, composition, and so forth) can be varied substantially, it should be possible to fine tune the properties of the composite. Moreover, the length scale of the microstructures of block copolymers is on the order of the characteristic length scale of the chains, ranging from 5 to 100 nm, which may make mesoporous materials with large pore sizes accessible. We investigated the sol-gel process of a mixture of two metal alkoxides, (3-glycidyloxypropyl)-trimethoxysilane, $(\text{CH}_3\text{O})_3\text{Si}(\text{CH}_2)_3\text{OCH}_2\text{CHCH}_2\text{O}$ (GLYMO), and aluminum *sec*-butoxide, $\text{Al}(\text{O}i\text{Bu})_3$, with poly(isoprene-*b*-ethylene-oxide) block copolymers (PI-*b*-PEO) (Fig. 1). GLYMO itself is an interesting hybrid material known to form thin-film coatings on polymers, thereby enhancing the abrasion resistance up to values of conventional glass (14). The block copolymer has two important features. First, the hydrolysis products of the metal alkoxides should preferentially swell the hydrophilic PEO block as a result of hydrogen bonding (as known, for example, from the synthesis of mesoporous molecular sieves with low-molecular weight nonionic surfactants) (6, 7). Second, the low glass transition temperature $T_g \approx 213$ K of the PI block introduces high mobility at ambient temperatures and should allow rapid formation of structures with long-range order even in the bulk.

Two PI-*b*-PEO block copolymers, referred to as PP3 and PP7, were synthesized by anionic polymerization using a recently described procedure (15). The molecular weights are nearly 10 kg mol^{-1} (PP3) and 34 kg mol^{-1} (PP7), and the polydispersity is low ($M_w/M_n \approx 1.05$, where M_w and M_n are the weight-average and number-average molecular weights, respectively). The volume fraction of the PEO block is $\sim 15\%$ in both cases. Their microdomain structure

was explored by small-angle x-ray scattering (SAXS) (Fig. 2). In a representative SAXS pattern obtained for PP3 at room temperature (Fig. 2A), the main peak is centered around a value for the scattering wave vector q corresponding to ~ 11.9 nm. There are at least two higher order reflections clearly visible at angular positions of $\sqrt{2}$ and $\sqrt{3}$ of the first-order maximum. This pattern is characteristic for spheres packed in a simple or body-centered cubic lattice, as expected for this volume fraction.

In a typical preparation of an organic-inorganic composite, 0.5 g of PI-*b*-PEO block copolymer was dissolved in a 1:1 mixture of CHCl_3 and tetrahydrofuran (5 weight % polymer); under moderate stirring, a pre-hydrolyzed solution of 80 mol % GLYMO and 20 mol % $\text{Al}(\text{O}i\text{Bu})_3$ (16) was added, and after 2 hours the mixture was transferred to a petri dish at 333 to 343 K. After subsequent evaporation of the organic solvents (~ 1 hour), the formation of the composite was accomplished by heat treatment at 403 K in vacuum for 45 min. A series of film samples with thicknesses of ~ 0.5 to 1 mm were prepared in this way by adding different amounts of the metal alkoxide solution to the same block copolymer. In the following, we focus on samples with 0.22 and 0.57 g of metal oxides in 0.5 g of PP3, denoted PP3/4 and PP3/10, respectively.

In the SAXS pattern of PP3/4 (Fig. 2B), the main peak is located at a q value corresponding to ~ 20.3 nm, and there are higher order reflections at angular positions of $\sqrt{4}$ and $\sqrt{7}$ of this first-order maximum. This spacing sequence is indicative of a hexagonal array of cylinders. For PP3/10 (Fig. 2C), the main peak is centered around a q value corresponding to ~ 19.6 nm, and two more reflections of higher order are clearly visible at integer multiples of this q value. Such a sequence is characteristic of an arrangement of lamellae.

To corroborate the assignment of these two SAXS patterns to a cylindrical and a lamellar morphology, respectively, we also examined the samples by transmission electron microscopy (TEM) (Fig. 3). The contrast in these micrographs arises from PI, stained with OsO_4 , and appearing black. The image of PP3/4 (Fig. 3A) clearly shows hexagonally packed cylinders in the two most typical projections. The TEM image of PP3/10 (Fig. 3B) exhibits lamellae. To determine whether the silica-type material is confined to one phase of the block copolymer, we used the recently developed method of elemental mapping (17). In Fig. 3C the silicon map of the same area depicted in Fig. 3B is shown; areas containing silicon appear bright in this image. Two conclusions can be drawn from Fig. 3C: (i) The inorganic silicon-rich phase

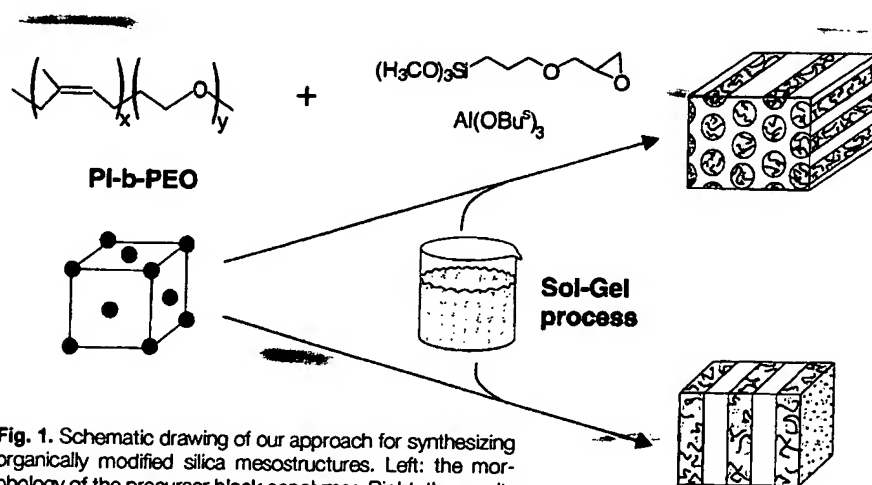


Fig. 1. Schematic drawing of our approach for synthesizing organically modified silica mesostructures. Left: the morphology of the precursor block copolymer. Right: the resulting morphologies after addition of various amounts of the metal alkoxides.

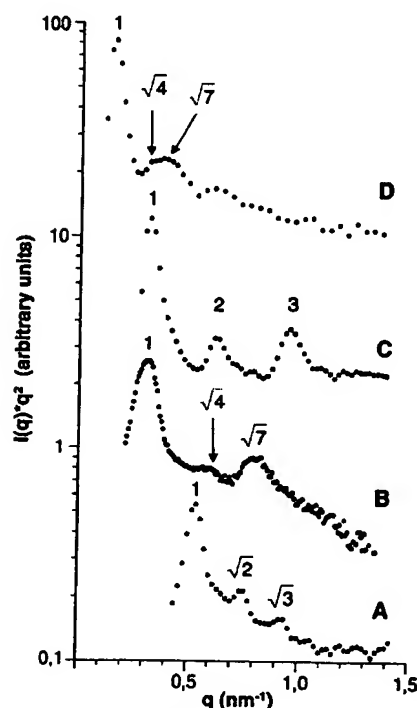


Fig. 2. Scattered intensities $I(q)$ as a function of scattering vector q for PP3 (A), PP3/4 (B), PP3/10 (C), and PP7/4 (D) at 295 K. Angular positions of higher order peaks with respect to the first-order maximum are indicated for each curve. The maxima with an angular position of $\sqrt{3}$, usually expected for a cylindrical morphology, are not well resolved in curves (B) and (D), probably because of the large width of the peaks. The patterns in (A), (C), and (D) were obtained with a Kratky compact camera (Anton Paar KG) equipped with a one-dimensional position-sensitive detector (M. Braun). The Ni-filtered $\text{Cu K}\alpha$ radiation ($\lambda = 0.154$ nm) was from a Siemens generator (Kristalloflex 710 H) operating at 35 kV and 30 mA. The pattern in (B) was obtained with a Rigaku Rotaflex x-ray source and a 2D area detector after integration over the azimuthal angle (see Fig. 4).

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has a lamellar morphology, and (ii) comparison of Fig. 3B and Fig. 3C shows that silicon is confined to the PEO phase of PP3. The

same spatial distribution can be shown by aluminum mapping, and similar results were obtained for the hexagonal phase (18).

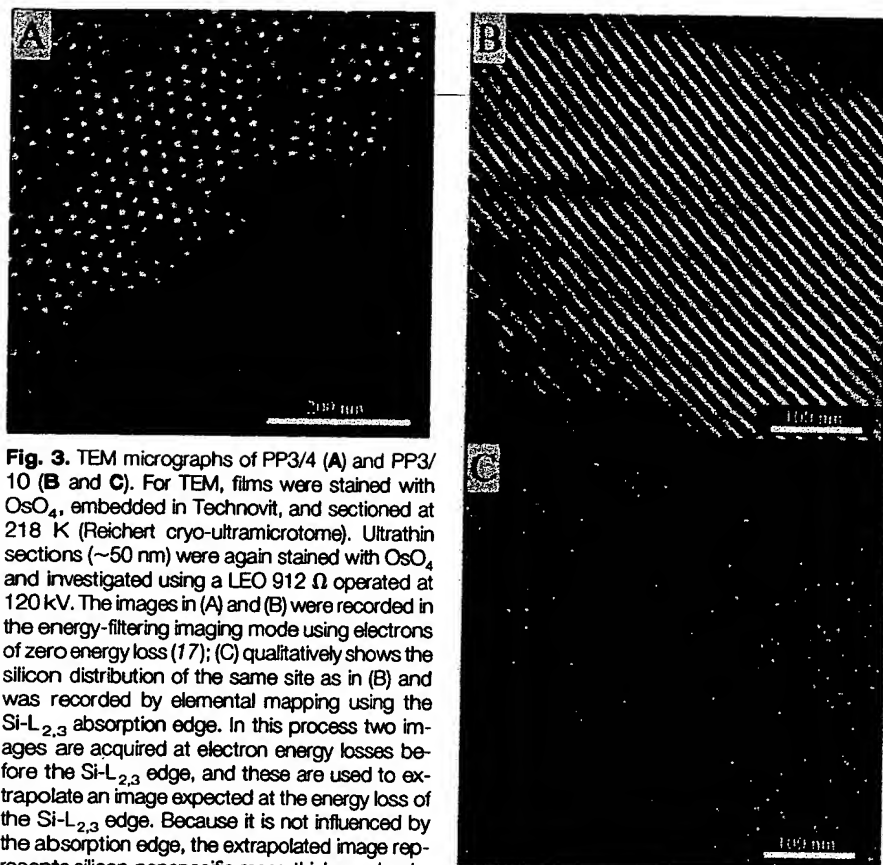


Fig. 3. TEM micrographs of PP3/4 (A) and PP3/10 (B and C). For TEM, films were stained with OsO_4 , embedded in Technovit, and sectioned at 218 K (Reichert cryo-ultramicrotome). Ultrathin sections (~ 50 nm) were again stained with OsO_4 and investigated using a LEO 912 Ω operated at 120 kV. The images in (A) and (B) were recorded in the energy-filtering imaging mode using electrons of zero energy loss (17); (C) qualitatively shows the silicon distribution of the same site as in (B) and was recorded by elemental mapping using the $\text{Si-L}_{2,3}$ absorption edge. In this process two images are acquired at electron energy losses before the $\text{Si-L}_{2,3}$ edge, and these are used to extrapolate an image expected at the energy loss of the $\text{Si-L}_{2,3}$ edge. Because it is not influenced by the absorption edge, the extrapolated image represents silicon-nonspecific mass-thickness background. This background is then subtracted from a third image acquired at the $\text{Si-L}_{2,3}$ edge, the difference image representing the pure distribution of silicon to the contrast (24). The smaller distance in the lamellar spacing in the TEM images relative to that indicated by the SAXS data is a result of contraction of the ultrathin sections normal to the plane of the lamellae, driven by free energy minimization (25).

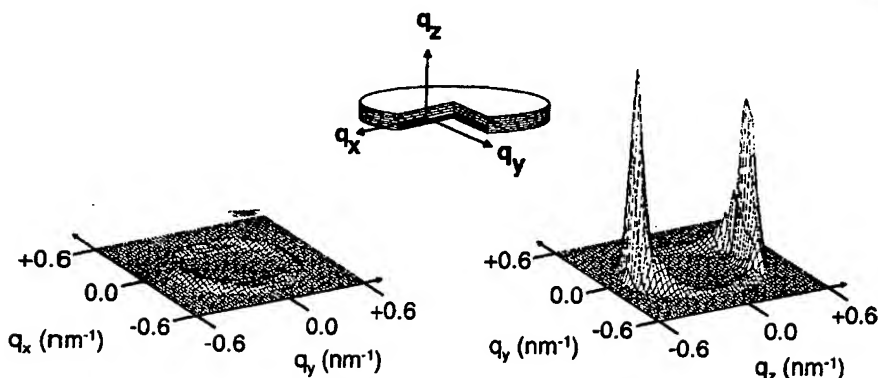


Fig. 4. Two-dimensional SAXS patterns of PP3/10 at 295 K for two different directions of the x-ray beam with respect to the sample coordinate frame, as schematically depicted in the inset. The order parameter (P_2) for the angular distribution of the lamellae normal with respect to the z-direction (film-normal), as obtained from the 2D SAXS pattern (21) on the right side, is 0.5. Patterns were obtained with a Rigaku Rotaflex x-ray source at 0.154 nm ($\text{Cu K}\alpha$). A three-pinhole collimator was used to generate a beam 1 mm in diameter. Scattering patterns were recorded on a 2D Siemens X-1000 area detector with a sample-to-detector distance of 130 cm.

The length scale of the morphologies in the present composites reflected from the SAXS patterns in Fig. 2, B and C, is ~ 20 nm, considerably longer than what is typically obtained for materials prepared from low-molecular weight surfactants. The spacings can be further increased by using higher molecular weight block copolymers. In composites prepared from PP7, spacings of ~ 40 nm were achieved (Fig. 2D). Because molecular weights of up to 10^2 to 10^3 kg mol^{-1} can be synthesized, fine-tuning of morphological parameters becomes possible.

The electron microscopy results (Fig. 3) suggest that the hydrophilic PEO block acts as an anchor for the metal alkoxide condensation products. More information about this effect was obtained from differential scanning calorimetry (DSC) (19). Although the T_g value of the PI block is unaffected, the melting behavior of the PEO block is markedly altered by the addition of inorganic material (18). For pure PP3, a melting point T_m was clearly detected at 310 K. For samples PP3/4 and PP3/10, however, the crystallization of the PEO block was suppressed. This is a well-known phenomenon in polymer blending, where the intimate mixing of a second polymer prevents PEO crystallization (20). Crystallization is only suppressed, however, if during the synthesis the organic solvents are evaporated at temperatures above the T_m of PEO. This suggests that the PEO chains and the hydrolysis products of the metal alkoxides mix well only above the T_m of the PEO block, and this state is frozen through condensation of the metal alkoxides.

Information about the inorganic connectivities can be gained by solid-state nuclear magnetic resonance. The condensation behavior of the present mixtures is similar to that of the pure metal alkoxides (21). Most of the silicon atoms are connected to two or three other metal atoms (silicon or aluminum) by oxygen bridges, thereby yielding a three-dimensional network. Nearly 40% of the aluminum is incorporated in this network as fourfold coordinated species. The residual aluminum is located in aluminum oxohydroxo complexes, $\text{AlO}_x(\text{OH})_y(\text{H}_2\text{O})_z$, as sixfold coordinated aluminum. In addition to the links on the inorganic side, the conversion of the epoxy group to oligoethyleneoxide derivatives leads to a higher network density.

Finally, we concentrate on orientational effects induced by the solvent-cast technique (22), which is part of the preparation procedure for our materials. Two-dimensional (2D) SAXS patterns for two different orientations of a film of lamellar sample PP3/10 with respect to the x-ray beam (Fig. 4) show that in the q_x - q_y plane (film plane) only a ring of small scattering intensity is

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observed, whereas in the q_y - q_z plane two strong and narrow scattering peaks along q_z are detected. This result is expected for lamellae oriented parallel to the film surface. It demonstrates that the solvent-cast technique is capable of inducing macroscopically aligned samples for the present lamellar silica-type mesostructures. Because the film thickness of these materials is considerable (~ 1 mm), surface-induced morphological transitions and related effects observed for very thin films (23) can be neglected.

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25. The decrease of the lamellar spacing is exclusively attributable to convex bending of the liquid-like lamellar surface. Two driving forces must be considered: surface tension (up to a critical ratio of lamella thickness to section thickness, the free lamella surface decreases upon bending) and entropy (partial relaxation of entropically unfavorable chain conformation (stretched chains in lamellae in bulk)). For details, see A. Du Chesne, thesis, University of Mainz (1993).
26. We thank H. W. Spiess for his strong support of this work, T. Thum-Albrecht for help in performing SAXS measurements and for fruitful discussions, and U. Pawelzik for performing the DSC measurements. M.T. thanks the Bundesministerium für Bildung, Wissenschaft, Forschung und Technologie (project 03 N 1009C0) for financial support.

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Promotion of the Cycling of Diet-Enhancing Nutrients by African Grazers

S. J. McNaughton,* F. F. Banyikwa, M. M. McNaughton

Experiments in Serengeti National Park, Tanzania, provide direct evidence that large, free-ranging mammalian grazers accelerate nutrient cycling in a natural ecosystem in a way that enhances their own carrying capacity. Both nitrogen and sodium were at considerably higher plant-available levels in soils of highly grazed sites than in soils of nearby areas where animal density is sparse. Fencing that uncoupled grazers and soils indicated that the animals promote nitrogen availability on soils of inherently similar fertility and select sites of higher sodium availability as well as enhancing that availability.

There is a growing recognition in ecology that organisms can modify their environments in ways beneficial to themselves, rather than inevitably causing environmental deterioration (1), and it is a maxim of grassland ecology that nutrient recycling by grazers contributes to plant regrowth potential (2). However, direct evidence of the effect of large wild mammals on nutrient recycling is meager (3), and studies in boreal forests (4) indicate that moose (*Alces alces*) browsing indirectly diminishes soil mineralization rate by shifting the composition of vegetation species to less palatable and less decomposable plants.

The distribution and abundance of large grazing mammals in Serengeti National Park, Tanzania, are influenced by the occurrence of nutritionally sufficient forages (5) and the spatiotemporal variation of vegetation productivity due to pronounced

geographic rainfall gradients and production seasonality (6). Grazers preferentially forage on swards enriched in minerals that are important in late-stage pregnancy, lactation, and the growth of young animals (5). There are two plausible explanations for this phenomenon: Animals forage on vegetation supported by soils of innately greater nutrient availability, or animal activities augment nutrient availability. Identification of the correct explanation has implications for conservation policy and management (through an understanding of the habitat requirements of endangered wild grazing mammals) and for ecological theory (by documenting how grazing mammals are mechanistically coupled with their habitats). Regional edaphic differences affect the mineral contents of forages and thereby influence seasonal movements of migratory grazers in the Serengeti, but no evidence of general soil differences was found in landscape-level studies of resident grazers (5), which are those that do not migrate but occupy discrete home ranges. Here we present evidence concerning the mechanisms associated with higher nutrient availability at sites preferred by resident grazers.

Mineralization of two elements—nitrogen (N), which is essential to both plants

S. J. McNaughton and M. M. McNaughton, Biological Research Laboratories, Syracuse University, Syracuse, NY 13244–1220, USA; and Serengeti Wildlife Research Centre, Post Office Box 661, Arusha, Tanzania. F. F. Banyikwa, Department of Botany, University of Dar es Salaam, Post Office Box 35060, Dar es Salaam, Tanzania; Biological Research Laboratories, Syracuse University, Syracuse, NY 13244–1220, USA; and Serengeti Wildlife Research Centre, Post Office Box 661, Arusha, Tanzania.

*To whom correspondence should be addressed: E-mail: sjmnaug@mailbox.syr.edu

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